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UNITED STATES ENVIRONMENTAL PROTECTION AGENCY WASHINGTON, D.C. 20460

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MAR 30 1992

MEMORANDUM

OFFICE OF PESTICIDES AND TOXIC SUBSTANCES

SUBJECT: Sulfuryl Fluoride (SF).

study in rats.

4 hour dermal vapor exposure

ID #1062719-00004

HED # 2-0109

Tox Chem No. 816A

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TO:

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Disinfectants Branch

Registration Division (H7505W)

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KB 124 92

Submitted Study:

"Sulfuryl Fluoride: Four-Hour Dermal Vapor Exposure in Fischer 344 Rats," by G. J. Bradley, T.D. Landry, J. E. Battjes and J. F. Quast of the Dow Chemical Company, Study ID# K-016399-036. MRID #417120-01.

The study was reviewed by Clement International Corporation, contractor for HED. The Clement review is attached to this memorandum.

Study Summary.

The purpose of this study was to determine any toxic effects from the absorption of SF vapor through the intact skin of rats while preventing the animals from breathing the SF to which their skins were exposed. Thus the animals were placed in a inhalation exposure chambers with their bodies inside the chambers and their heads outside in a reversed head only exposure position. The bodies were then exposed for 4 hours to SF at 9599 ppm SF. No adverse effects were observed from these exposures.

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Conclusions.

Based on the results of this study, the dermal vapor LC50 of SF in rats is greater than 9599 ppm (40.3 mg/L). No TOXICITY CATEGORY can be assigned to this study because we have not corresponding category. Based on the methods and observations used, the study is ACCEPTABLE.

Comments:

This study is not typical of guideline specifications; however it met its objectives of assessing the possible toxicity from dermal exposures to high concentrations of SF. The vapor exposures were well performed and characterized. Whole animal toxicity of sevaluation along with possible skin and internal organ pathology and toxicity of sevaluation along with possible skin and internal

Note; A one-liner summary for this study was attached with the SF/RED toxicology summary prepared for FIFRA 1988 review of SF.

DOC9A0123 FINAL

DATA EVALUATION REPORT

SULFURYL FLUORIDE

Study Type: Dermal Vapor Exposure in Rats

Study Title: Sulfuryl Fluoride: Four-Hour Dermal Vapor Exposure in Fischer

344 Rats

Prepared for:

Health Effects Division Office of Pesticide Programs Environmental Protection Agency 1921 Jefferson Davis Highway Arlington, VA 22202

Prepared by:

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Principal Reviewer

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2/11/92 Date

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Contract Number: 68D10075
Work Assignment Number: 1-35

Clement Number: 91-127

Project Officer: James E. Scott

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EPA Reviewer: Stanley Gross, Ph.D.

Review Section II, Toxicology Branch I/HED

EPA Section Head: Joycelyn Stewart, Ph.D. Review Section II, Toxicology Branch I/HED Signature: Date:

Signature:

Date:

DATA EVALUATION REPORT

STUDY TYPE: Modified Guideline Series 81-2: Acute Dermal Toxicity Study in Rats

EPA IDENTIFICATION NUMBERS

Tox. Chem. Number:

MRID Number: 417120-01

TEST MATERIAL: Sulfuryl fluoride

SYNONYMS: VIKANE gas fumigant

SPONSOR: DowElanco, Midland, MI

STUDY NUMBER: K-016399-036

TESTING FACILITY: The Toxicology Research Laboratory, Health and Environmental Sciences, The Dow Chemical Company, Midland, MI

TITLE OF REPORT: Sulfuryl Fluoride: Four-Hour Dermal Vapor Exposure in

Fischer 344 Rats

AUTHOR: G.J. Bradley, T.D. Landry, J.E. Battjes, and J.F. Quast

STUDY COMPLETED: November 16, 1990

CONCLUSIONS: The dermal vapor LC₅₀ for sulfuryl fluoride in male and female rats is greater than 9599 ppm (40,315 mg/m³).

ONE-LINER: The No-Observed-Effect Level (NOEL) for dermal exposure to sulfuryl fluoride vapor in rats is 9599 ppm (40,315 mg/m³).

CORE CLASSIFICATION: This study is acceptable (modified guideline series 81-2).

TOXICITY CATEGORY: Not applicable

A. MATERIALS

1. Test Material

Compound: Sulfuryl fluoride (SO₂F₂)

Purity of material: 99.67%

Physical description: Colorless gas

Lot no.: 880329 752

Storage conditions: Not reported

2. Controls

Materials: None Animals: None

3. Test Animals

Species: Rats

Strain: Fischer 344

Source: Charles River Breeding Laboratories, Inc., Kingston, NY

Receipt date: Not reported

Sex: Male and female

Numbers: 10 males, 10 females

Housing: Double

Age: 6-8 weeks at time of exposure

Weight: Preexposure: 101.2-122.9 g (males), 90.5-126.8 g (females)

Feeding: Feed and water ad libitum

Assignment: Computer-generated randomization

4. Exposure

Limit test:

Route of administration: Dermal vapor

Dose level: 987 ppm (5 males)

1013 ppm (5 females)

9599 ppm (5 animals/sex)

B. TEST PERFORMANCE

Prior to exposure, the dorsal surface of each rat was shaved with electric clippers.

Exposure Concentration

Groups of five male or five female rats were dermally exposed (body only) to 987 ppm and 1013 ppm of sulfuryl fluoride, respectively, for a single 4-hour period. Since no effects were noted from this exposure, another group of five rats/sex/dose group were dermally exposed to 9599 ppm for

4 hours. This concentration (9599 ppm) was reported to be approximately 10-fold greater than the whole-body inhalation ${\rm LC}_{50}$.

Chamber

A stainless steel and glass, 157-liter, Rochester-type chamber (50 cm x 50 cm x 50 cm) was used. The chamber was modified so that the heads of the rats protruded through an elastic dental dam, which served as a barrier between the test material and the breathing air for the rats. A ventilated enclosure (*30 liters/minute) surrounded the protruding heads to allow monitoring of the air the animals were breathing. Chamber air was controlled by a system designed to maintain temperature and humidity at 22°C and 50%, respectively. Air flow through the chamber was maintained at approximately 30 liters per minute. Rats were dermally exposed to vapors of the tesc material and were housed individually to minimize crowding during the exposure period. A diagram of the exposure chamber was provided.

Vapor Generation

Sulfuryl fluoride vapors were generated by metering from a gas sampling bag made of SARAN® resin with an FMI pump (Fluid Metering Inc., Oyster Bay, NY) to a J-tube. Compressed air was mixed with the sulfuryl fluoride gas in the glass J-tube assembly to attain the desired chamber concentration.

Chamber Monitoring

Air flow through the chamber was determined with a precalibrated manometer prior to study initiation. Temperature, relative humidity, and air flow values were recorded every 30 minutes during the 4-hour exposure. The analytical concentration of sulfuryl fluoride in the chamber was measured at least once per hour with a MIRAN 1A infrared spectrophotometer. The breathing area of the rats was checked at least once per hour to determine if there was significant inhalation exposure to sulfuryl fluoride (minimum analytical sensitivity was <20 ppm). A distribution check of test material in the breathing zone was performed prior to exposure of the animals to the test material. Results showed that vapor concentrations in the four sampling ports ranged from 102% to 104% of the standard.

Observations

Animals were weighed and examined prior to test material exposure (day 1). All rats were weighed on test days 2, 4, 8, 11, and 15 during the 2-week postexposure period. Animals were observed during exposure and daily during the 2-week postexposure period. Fur, eyes, mucous membranes, and respiration were examined. Behavior pattern and nervous system activity

¹Miller, R.R., Calhoun, L.L., Keyes, D.G., and Kociba, R.J. 1980a. Sulfuryl Fluoride (VIKANE® fumigant): An LC₅₀ Determination. Report of the Dow Chemical Company Toxicology Research Laboratory, Midland, MI. were also assessed by observing for tremors, convulsions, salivation, lacrimation, diarrhea, lethargy, and other signs of altered central nervous system function.

All rats were subjected to gross necropsy on test day 15; brain and samples of clipped and unclipped skin were saved. Multiple sections of brain and clipped and unclipped skin from each animal in the 9599-ppm group were processed by routine histologic procedures. No gross pathologic observations were noted in any rats; therefore, no other tissues were microscopically examined.

C. RESULTS AND STUDY AUTHORS' CONCLUSIONS

Tables were presented for chamber atmosphere conditions during exposure, clinical observations, body weights, gross pathology, and histopathologic observations. Findings were as follows:

Analytical Determinations

Test atmospheres in the exposure chamber were reported to be 987 ± 21 , 1013 ± 9 , and 9599 ± 142 ppm. Chamber temperature (23-24°C) and relative humidity (33-39%) were generally comparable for each exposure.

Animal Observations

Since no compound-related effects were noted in the five male and five female rats exposed to 987 ppm and 1013 ppm, respectively, an additional group of five rats/sex/dose were dermally exposed to 9599 ppm for 4 hours.

Animals in all groups survived the 4-hour exposure and the 14-day postexposure period. The majority of rats in all exposure groups exhibited chromodacryorrhea and fecal soiling. The study authors reported that these findings were related to the method of restraint during the exposure period.

Average body weights decreased slightly (<3%) from pretreatment values immediately following exposure to all concentrations of the test substance. The study author reported that this weight loss was not treatment related. Similar weight losses occurred during a preliminary assessment performed on rats in the chamber prior to the study. By day 4, all animals exceeded their pretreatment weights and continued to gain weight throughout the remainder of the study.

Gross pathological examinations of all animals were normal. Histopathologic examinations of brain and skin samples taken from animals exposed to 9599 ppm revealed no treatment-related lesions.

D. REVIEWERS' COMMENTS

Based on the results of this study, the dermal vapor LC_{50} of sulfuryl fluoride in rats is greater than 9599 ppm. No toxicity category was

assigned to this study because there is no corresponding category. Based on the observations and methods used, this study is acceptable.

One-Liner

Based on the results of this study, the NOEL for dermal exposure to sulfuryl fluoride vapor in rats is 9599 ppm (40,315 mg/m 3).

E. QUALITY ASSURANCE MEASURE

A signed Quality Assurance Statement, dated 11/16/90, was presented. A Good Laboratory Practice compliance statement was included.

F. CBI APPENDIX

CBI Materials and Methods, pp. 8-13.

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Identity of product impurities.	
Description of the product manufacturing proces	ss.
Description of quality control procedures.	
Identity of the source of product ingredients.	
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A draft product label.	•
The product confidential statement of formula.	
Information about a pending registration actio	n.
FIFRA registration data.	
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